

1 **Title:** Respiration rates and active carbon flux of mesopelagic fishes (Family Myctophidae) in the  
2 Scotia Sea, Southern Ocean

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6 **Running page head:** Myctophid respiration in the Southern Ocean

7 **Keywords:** Myctophid, Respiration, Lantern fish, Carbon, Active Flux, Southern Ocean

8 **Abstract**

9 Mesopelagic fish have recently been highlighted as an important, but poorly studied component of  
10 marine ecosystems, particularly regarding their role in the marine pelagic food webs and  
11 biogeochemical cycles. Myctophids (Family Myctophidae) are one of the most biomass-dominant  
12 groups of mesopelagic fishes, and their large vertical migrations provide means of rapid transfer of  
13 carbon to the deep ocean where it can be sequestered for centuries or more. In this study, we  
14 develop a simple regression for the respiration rate of myctophid fish using literature-based wet  
15 mass and habitat temperature data. We apply this regression to net haul data collected across the  
16 Scotia-Weddell sector of the Southern Ocean to estimate respiration rates of the biomass-dominant  
17 myctophid species. *Electrona carlsbergi*, *Electrona antarctica* and *Gymnoscopelus braueri* made a  
18 high contribution (up to 85%) to total myctophid respiration. Despite the lower temperatures of the  
19 southern Scotia Sea (-1.46 to 0.95 °C), total respiration here was as high (reaching 1.1 mg C m<sup>-2</sup> d<sup>-1</sup>)  
20 as in the warmer waters of the mid and northern Scotia Sea. The maximum respiratory carbon flux  
21 of the vertically migrating community was 0.05-0.28 mg C m<sup>-2</sup> d<sup>-1</sup>, equivalent to up to 47% of the  
22 gravitational particulate organic carbon flux in some parts of the Scotia-Weddell region. Our study  
23 provides the first baseline estimates of respiration rates and carbon flux of myctophids in the  
24 Southern Ocean. However, direct measurements of myctophid respiration, and of mesopelagic fish  
25 generally, are needed to constrain these estimates further and incorporate these fluxes into carbon  
26 budgets.

## 27 **Introduction**

28 The biological uptake and cycling of carbon in the ocean are tightly coupled to atmospheric  
29 levels of carbon dioxide (CO<sub>2</sub>) (Sabine et al. 2004). Primary production in the surface ocean  
30 drives the uptake of CO<sub>2</sub>, but it only begins to be sequestered once it is transferred below  
31 the mixed layer and is no longer in contact with the atmosphere (Primeau 2005). Species  
32 that migrate vertically in the water column can actively transfer carbon to the deep ocean  
33 through excretion, defecation, mortality and respiration (Longhurst et al. 1990, Zhang &  
34 Dam 1997, Steinberg et al. 2000, Turner 2002, Steinberg & Landry 2017, and references  
35 therein). This has been studied greatly in marine zooplankton (e.g. Zhang & Dam 1997,  
36 Steinberg et al. 2000, Hernández-León et al. 2001, Packard & Gómez 2013), however, there  
37 have been few studies examining active transport in migratory fish, particularly mesopelagic  
38 fish (e.g. Hidaka et al. 2001; Davison et al. 2013; Hudson et al. 2014; Ariza et al. 2015), which  
39 are difficult to sample effectively in remote open ocean regions.

40 Recently, the importance of including mesopelagic fish in ocean carbon budgets has been  
41 highlighted (Anderson et al. 2018). They are one of the components of marine ecosystems  
42 that we know least about (St John et al. 2016), yet they are highly motile and many species  
43 migrate vertically, feeding at the surface during the night, but migrating to the mesopelagic  
44 and bathypelagic zones during the day where they continue to respire. Previous studies  
45 have found the respiratory carbon flux of migratory fishes to be equivalent to up to 26% of  
46 the gravitational particulate organic carbon (POC) flux (Hidaka et al. 2001, Hudson et al.  
47 2014, Ariza et al. 2015). In addition, their gut passage times are much slower than  
48 zooplankton (Ariza et al. 2015), and thus faecal pellets are more likely to be released in the  
49 deep ocean following night-time feeding at the surface.

50 Lantern fish (Family Myctophidae, here after myctophids) are the most common  
51 mesopelagic fish in most of the World's oceans (Catul et al. 2011), and are known to make  
52 large vertical migrations (Pakhomov et al. 1996). In the mesopelagic and bathypelagic zones  
53 of the Southern Ocean, they are the dominant fish family in terms of species richness,  
54 abundance, and biomass (Duhamel et al. 2014), and are important in the pelagic ecosystem  
55 in this region (Murphy et al. 2007). Yet there have been no studies attempting to quantify  
56 the contribution of myctophid species in the Southern Ocean to active carbon fluxes.  
57 Indeed, the role of mesopelagic and bathypelagic fish communities in biogeochemical  
58 cycling and carbon transfer to depth is one requiring urgent research, both regionally and  
59 globally (Trueman et al. 2014).

60 The respiration rates of myctophid fish are not easy to measure directly, due to difficulties  
61 in obtaining live, healthy specimens from the mesopelagic zone, and our inability to  
62 successfully incubate them under stress-free conditions. Therefore, previous studies (Hidaka  
63 et al. 2001, Hudson et al. 2014) examining myctophid respiration have either utilised the  
64 relationship between biomass and respiration established by the historical study of Donnelly  
65 and Torres (1988), or used general allometric relationships between mass and metabolic  
66 rate for other fish (Davison et al. 2013). An exception is Ariza et al. (2015), who made direct  
67 measurements of electron transport system (ETS) activity in order to estimate respiration. A  
68 number of large compilations of respiration data have been made, defining regressions  
69 between the biomass of marine organisms and their respiration (e.g. Ikeda et al. 2001; Ikeda  
70 2016), yet none of these were specific to myctophid species. There can be significant  
71 variation in the resting metabolism, and hence, routine respiration, of different taxonomic

72 groups (Clarke & Johnston 1999). Therefore, generalised regressions for pelagic marine  
73 fishes (Ikeda 2016) may not provide the most accurate estimate of myctophid respiration.  
74 In this study, we compile previous estimates of myctophid respiration from the literature to  
75 define a simple regression to calculate myctophid respiration from wet mass and habitat  
76 temperature. We then utilise net haul data, collected as part of the most comprehensive  
77 mesopelagic fish survey in the Southern Ocean to date, to examine myctophid respiration in  
78 the Scotia Sea, one of the most productive regions of the Southern Ocean. In this way, we  
79 start to quantify their importance in the active transfer of carbon to depth.

## 80 **Methods**

### 81 *Myctophid distribution and abundance*

82 Detailed surveys for mesopelagic fish were conducted in the Scotia Sea as part of the British  
83 Antarctic Survey's Discovery 2010 programme, as has been previously described in Collins et  
84 al. (2012). Briefly, this involved deployment of an opening and closing 25 m<sup>2</sup> rectangular  
85 mid-water trawl net (RMT25, minimum 4 mm mesh; Piatkowski et al. 1994) along a transect  
86 spanning the entire Scotia Sea between the Antarctic Polar Front (APF) and the sea ice zone  
87 (SIZ) during three cruises; in November 2006 (cruise JR161, Austral spring), January 2008  
88 (cruise JR177, Austral summer), and March 2009 (cruise JR200, Austral autumn). Depth-  
89 stratified net hauls were carried out at six stations that encompassed the main water  
90 masses and frontal zones of the region: Polar Front (PF), Southern Scotia Sea (SSS), Mid  
91 Scotia Sea (MSS), Western Scotia Sea (WSS), Northern Scotia Sea (NSS), and Georgia Basin  
92 (GB). At each station, an RMT25 was deployed at the depth zones: 0-200, 200-400, 400-700,  
93 and 700-1000 m. The depth and ambient temperature of the nets were logged using a

94 custom-built net monitoring system. The temperature sensor (SBE-3) was factory calibrated  
95 prior to the surveys and was accurate to  $\sim 0.001$  °C. Net hauls were repeated during the day  
96 and night in spring and summer, but only during the night-time in autumn. All fish caught  
97 were sorted onboard, identified to the lowest taxonomic level, measured to the nearest mm  
98 using standard length (SL) and the wet mass (WM) measured to the nearest 0.01 g using a  
99 motion compensated balance. General patterns in community structure of these  
100 mesopelagic fish can be found in Collins et al. (2012) and information on species-specific  
101 biomass, abundance and population dynamics of the main myctophids is detailed in  
102 Saunders et al. (2014, 2015a, b).

103 For 39% of data records (23, 9, and 97% for JR161, JR177 and JR200 cruises, respectively),  
104 the WM was not measured and only the standard length of the fish was recorded. In these  
105 instances, we used length-mass regressions from the long-term records held at the British  
106 Antarctic Survey (unpubl. data, supplementary Table S1). Where possible, these were  
107 species-specific, or else genus-specific for the rarer species. Overall, individual fish WM  
108 ranged from 0.03 to 78.34 g (mean 4.38 g).

### 109 *Myctophid respiration regression*

110 To calculate the total myctophid respiration at each of the sites sampled, we developed a  
111 regression based on literature measurements of myctophid respiration. A search of the  
112 literature was carried out to identify studies in which the respiration rate of myctophids was  
113 measured, and the temperature and body mass (in terms of wet mass (WM), dry mass (DM)  
114 or carbon (C)) were also recorded. We identified 5 such studies (Torres et al. 1979, Donnelly  
115 & Torres 1988, Torres & Somero 1988, Ikeda 1989, Ariza et al. 2015), giving a total of 74  
116 data points from which we could base our regression analysis (Table 1).

117 Torres et al. (1979), Donnelly and Torres (1988) and Torres and Somero (1988) measured  
118 the routine respiration (i.e. under conditions of normal activity) via incubations at  
119 temperatures experienced *in situ*. Both Ikeda (1989) and Ariza et al. (2015) measured the  
120 capacity of the respiratory ETS, converting this potential respiration to the actual respiration  
121 via experimentally determined ratios. Where possible, we have compiled respiration and  
122 WM data for individual fish. However, in instances where the individual-specific data were  
123 unavailable, we took either the given mean WM and respiration, or in the case of Torres et  
124 al. (1979), the calculated mean WM for the given range.

125 As the aim was to develop a regression that could readily be applied to fish catch data  
126 collected in the field, we chose to develop an equation for the WM specific respiration rate  
127 ( $R_{WM}$ , in  $\mu\text{L O}_2 \text{ mg WM}^{-1} \text{ h}^{-1}$ ) from fish WM (in mg) and ambient temperature ( $T$ , °C). Based  
128 on relationships previously established between biomass and respiration (Kjørboe & Hirst  
129 2014, Ikeda 2016), we define a simple regression model.

130

$$131 \quad \ln(R_{WM}) = a_0 + a_1 \times \ln(WM) + a_2 \times T \quad (1)$$

132

133 Here  $a_0$ ,  $a_1$ , and  $a_2$  are regression coefficients. Regression analysis was carried out using a  
134 regression fitting model for multiple predictors and a response, where data were  
135 continuous and no interactions terms were allowed. Wet mass and respiration data were  
136 transformed to the natural log prior to fitting the regression. Fitting was performed using  
137 the ordinary least squares method in Minitab 18 (version 18.1). To assess the uncertainty  
138 surrounding our calculated regression coefficients, we applied bootstrapping. For this

139 procedure, we randomly sampled (with replacement) from all 74 literature-based data  
140 points on myctophid fish respiration to generate 100 simulated dataset. We then calculated  
141 the regression coefficients (as above) for each of these data sets and in this way, estimated  
142 bootstrapped confidence intervals (standard error) for each coefficient over the 100  
143 simulations.

#### 144 *Total respiration*

145 We combine the results of our regression model with the Discovery 2010 survey data to  
146 calculate the respiration rate for each individual fish ( $R_{IND}$ ,  $\mu\text{L O}_2 \text{ ind.}^{-1} \text{ h}^{-1}$ ) in a particular net  
147 haul. The total respiration  $R_{TOT}$  ( $\mu\text{L O}_2 \text{ m}^{-3} \text{ h}^{-1}$ ) for each net haul was then calculated by  
148 standardising to the volume filtered by the net ( $V$ ,  $\text{m}^3$ ), and summing for all myctophid  
149 individuals captured in that haul.

150

$$151 \quad R_{TOT} = \sum \frac{R_{IND}}{V} \quad (2)$$

152

153 This was then converted to units of carbon per day ( $R_{TOT,C}$ ,  $\text{mg C m}^{-3} \text{ d}^{-1}$ ) using a respiratory  
154 quotient (RQ) of 0.90 for fishes (Brett & Groves 1979, Ariza et al. 2015) and the  
155 stoichiometric relationship between carbon and oxygen ( $22.4 \text{ L O}_2 = 12 \text{ g carbon}$ ). For each  
156 cruise, at each station, the mean  $R_{TOT}$  of any replicate hauls was calculated for each depth  
157 horizon. This was computed for day and night hauls separately. Only the night-time data  
158 were used for inter-station and inter-species comparisons of total respiration due to the  
159 inherent problem of daytime net avoidance by myctophid fish (Pakhomov et al. 1996,  
160 Collins et al. 2012) (see below).

161 *Maximum respiratory flux*

162 Many myctophid species are active migrators moving to the euphotic zone at night and  
163 returning to depth during the day, fluxing carbon to depth in the process. The maximum  
164 respiratory flux (below 200 m) of the migrant myctophid community was calculated by  
165 comparing  $R_{TOT,C}$  in the 0-200 m depth strata during the day and night (i.e. we subtract the  
166 total respiration of the resident community, the day-time respiration ( $R_d$ ), from the  
167 respiration of the night-time community ( $R_n$ )). Weather and net failure constraints during  
168 the Discovery 2010 cruises resulted in these calculations being possible for four stations,  
169 JR161 WSS and NSS, and JR177 MSS and GB. Our respiration calculations for the 0-200 m  
170 depth horizon are based on the ambient temperature over this depth range, but migrating  
171 individuals will experience different temperatures at depth. Therefore, to estimate the  
172 respiration of the migrating community at depth, we recalculated respiration rates using the  
173 mean temperature at depths of 400-1000 m. Finally, the maximum daily downward flux of  
174 respiratory carbon below 200 m by myctophid migrants ( $R_m$ ) was estimated based on the  
175 number of daylight hours ( $h$ ) at each station over the period of the research cruise (mean of  
176 the maximum and minimum daylight length).

177

178 
$$R_m = (R_n - R_d) \times \frac{h}{24} \quad (3)$$

179

180 We stress here that these calculations represent the maximum respiratory carbon flux. This  
181 is due to the issue of daytime net avoidance (Collins et al. 2012, Fielding et al. 2012). To  
182 investigate this uncertainty, we conducted a sensitivity analysis by recalculating day-time

183 respiration assuming catch efficiencies of 14%, 25% and 50%, and used these revised values  
184 for sensitivity analysis of the respiratory carbon flux of the migrant myctophid community.

## 185 **Results**

### 186 *Myctophid respiration regression*

187 The compiled respiration dataset comprised of myctophids (18 species, plus 23 individuals  
188 identified to the genus *Myctophum*) of WM ranging from 0.026 – 19.2 g, and experimental  
189 temperatures from 0.5 to 27 °C (Figure 1). The respiration rates (mass specific) decrease  
190 with increasing WM and increase with increasing temperature (Figure 1).

191 Regression analysis of the collated data reveals the following regression for mass specific  
192 respiration ( $R_{WM}$ ) of myctophid fishes ( $n=74$ , adjusted  $R^2=0.85$ ), with standard error of  
193 coefficients shown in brackets:

194

$$195 \quad \ln(R_{WM}) = -1.315 (\pm 0.468) - 0.2665 (\pm 0.0516) \times \ln(WM) + 0.0848 (\pm 0.0108) \times T \quad (4)$$

196

197 The standard errors calculated from our bootstrap analysis were 0.0368, 0.0040 and 0.0010  
198 for  $a_0$ ,  $a_1$ , and  $a_2$  respectively.  $R_{WM}$  increases with increasing temperatures (supplementary  
199 Figure S1) and decreases with increasing wet mass (supplementary Figure S2).

### 200 *Myctophid respiration: Seasonal changes*

201 Total respiration was calculated for each haul of the Discovery 2010 cruises, highlighting  
202 both latitudinal and seasonal patterns. We present the seasonal change in total myctophid

203 respiration for the NSS, MSS and SSS stations (Figure 2) as these are the stations where we  
204 have data from all four depth horizons on all three cruises. Night-time only data is examined  
205 to avoid bias by net avoidance during the day. Total respiration (integrated from 0-1000 m  
206 depth) was highest at SSS in autumn ( $1.0 \text{ mg C m}^{-2} \text{ d}^{-1}$ ), with the lowest rates occurring at  
207 NSS in autumn ( $0.4 \text{ mg C m}^{-2} \text{ d}^{-1}$ ). Whereas total respiration increased from spring to  
208 autumn at SSS, the opposite pattern was observed at NSS. Total respiration peaked at  $1.0$   
209  $\text{mg C m}^{-2} \text{ d}^{-1}$  in summer at MSS.

210 Seasonal differences were also apparent in the species making the dominant contribution to  
211 the total respiration (Figures 3-5). At NSS (Figure 3), *Electrona carlsbergi* accounted for 51%  
212 of the total respiration in spring. As the season progressed at NSS, the total respiration  
213 decreased for all species except *Electrona antarctica*, which peaked in summer, and the  
214 contribution to total respiration was much more equal across the different species.

215 At MSS (Figure 4), the highest total respiration was also due to *E. carlsbergi* but, in this case,  
216 this occurred in the summer, contributing 43% to the total respiration. *E. antarctica* also  
217 made a strong contribution (26%) to total respiration at MSS in summer. In both the spring  
218 and autumn, *Gymnoscopelus braueri* dominated the total respiration (38 and 33%  
219 respectively). At SSS (Figure 5), *E. antarctica* and *G. braueri* were the dominant species in  
220 terms of total respiration, with *G. braueri* dominating in spring (39%) and *E. antarctica*  
221 dominating in summer (47%) and autumn (45%).

#### 222 *Myctophid respiration: depth stratified, day – night comparisons*

223 There are four sites where we have complete day and night data for all four depth horizons  
224 (Figure 6), WSS and NSS in the spring, and GB and MSS in the summer. Total respiration was

225 highest at night-time, possibly because this was when more fish were caught, however, the  
226 potential net avoidance during the day makes it difficult to ascertain exact migration  
227 patterns. In the summer, *E. antarctica* dominated the total depth integrated respiration  
228 during both the day and night at MSS, however, during the day, respiration was highest in  
229 the 0-200 and 401-700 m depth horizons (0.0007 and 0.0006 mg C m<sup>-3</sup> d<sup>-1</sup> respectively)  
230 whereas, at night, respiration of *E. antarctica* was highest (0.0009 mg C m<sup>-3</sup> d<sup>-1</sup>) in the 701-  
231 1000 m depth range. Generally there was a decline in the total respiration with depth during  
232 the night, and an increase with depth during the day.

233 Although a particular species may dominate the total depth integrated respiration, this may  
234 be confined to particular depth horizons (Figure 6). For example, *E. carlsbergi* appears to  
235 contribute markedly to the total respiration at NSS in spring and MSS in summer (Figure 6),  
236 but our data suggest that its contribution is limited to the upper 400 m of the water column.  
237 Conversely, in the summer, both *E. antarctica* and *G. braueri* were important contributors to  
238 the myctophid respiration at all depth horizons during the day and night, with possible net  
239 avoidance or migration out of the top 200 m during the day.

#### 240 *Maximum respiratory flux*

241 Of the four sites where data were sufficient, the maximum respiratory flux of carbon below  
242 200 m by the migrant myctophid community was highest at NSS in the spring (0.28 mg C m<sup>-2</sup>  
243 d<sup>-1</sup>). The maximum respiratory carbon flux at GB in summer was lower (0.13 mg C m<sup>-2</sup> d<sup>-1</sup>),  
244 with the lowest flux of 0.05 mg C m<sup>-2</sup> d<sup>-1</sup> at WSS in spring.

245 As net sampling of nekton is not 100% efficient, with net avoidance being a particular  
246 problem during the daytime, we conducted a sensitivity analysis to examine how this alters

247 our calculations of the respiratory carbon flux. Studies have found net capture efficiencies of  
248 ~14% for net mouth areas between 5 and 105 m<sup>2</sup> (Koslow et al. 1997, Davison 2011). We  
249 take this as a lower bound estimate for our sensitivity analysis, recalculating the respiratory  
250 carbon flux based on day-time capture efficiencies of 14, 25, and 50% (Table 3).

251 Our sensitivity analysis highlights that these uncertainties in catch efficiency present  
252 problems for accurately incorporating these fluxes into mesopelagic carbon budgets. In two  
253 instances (JR161 WSS and JR177 GB), the respiratory flux assuming 14% day-time capture  
254 efficiency results in slightly negative estimates of respiratory carbon flux. However, it is also  
255 likely that there is also some net avoidance at night-time which we have not attempted to  
256 account for here due to unknown catch efficiencies.

## 257 **Discussion**

### 258 *Catch efficiency*

259 Considering the lack of data on mesopelagic fish respiration, and difficulty of obtaining such  
260 data, we attempt here to estimate respiration of the dominant fish, myctophids, in the  
261 Southern Ocean, based on biomass and temperature data. In this way we can start to assess  
262 the importance of mesopelagic fish in the Southern Ocean carbon budget. Although our  
263 calculations are based on a dedicated survey programme, spanning multiple regions and  
264 seasons, the biomass data are from net hauls and hence suffer the problems of net  
265 avoidance and catch efficiency.

266 The sampling of fish, and miconekton generally, via nets is fraught with the loss of  
267 individuals due to both net avoidance by large, fast swimmers during the day, and the loss  
268 of smaller animals through the mesh of the net. The capture efficiency is related to the net

269 design as well as to the size and swimming ability of micronekton (Gartner et al. 1989, Itaya  
270 et al. 2007), it is therefore not possible to apply a single correction factor. Acoustic  
271 estimates of biomass are generally greater than those from net trawls (e.g. Koslow et al.  
272 1997; Kaartvedt et al. 2012; Davison et al. 2015a), but acoustic estimates of mesopelagic  
273 fish biomass also present several challenges and require thorough ground-truthing (Davison,  
274 Koslow, et al. 2015). The sensitivity analysis that we conducted (Table 3) increases the range  
275 of our estimates of respiratory active flux, highlighting the need for new developments in  
276 acoustic techniques to improve myctophid abundance estimation that will further constrain  
277 estimates of respiratory flux by mesopelagic fish.

#### 278 *Myctophid respiration regression*

279 Analysis of the myctophid data we collated shows the expected trends of increasing mass  
280 specific respiration with increasing temperature and, decreasing mass specific respiration  
281 with increasing WM, as have been found by previous respiration studies (Winberg 1956,  
282 Clarke & Johnston 1999, Ikeda 2016). The aim of this study is not to examine the theory  
283 behind the success of various predictors, but to develop a simple equation to make first  
284 order estimates of the respiration of myctophid fishes. Our regression therefore uses  
285 parameters that are easily measurable in the field, T and WM.

286 Although our respiration regression is predominantly driven by abundance and WM, we do  
287 not see the same patterns for respiration as have been shown for abundance for the  
288 Discovery 2010 data. The calculated respiration depends on not only the total biomass, but  
289 also on the contribution of different sized fishes to the total biomass. For example, we  
290 would calculate much higher mass specific respiration (and lower total respiration) for a site  
291 with large numbers of small sized individuals, compared to a site with the same biomass but

292 comprised of fewer numbers of larger individuals. As the *in situ* temperature of the data  
293 used from the Discovery 2010 cruises had a small range of -1.46 – 3.31 °C (based on mean  
294 net haul temperatures), temperature plays a smaller role in the differences in respiration  
295 between stations.

296 The regression we have developed is based on a relatively small number of studies ( $n=5$ )  
297 and data points ( $n=74$ ), each of which is associated with methodological weaknesses. Torres  
298 et al. (1979), Donnelly and Torres (1988) and Torres and Somero (1988) conducted  
299 incubations on live fish to measure respiration. These incubation-based measurements can  
300 introduce errors due to stress during net capture and incubation, starvation and bacterial  
301 growth. This is particularly true in highly motile myctophid fish that migrate in the water  
302 column. Although the ETS method adopted by Ikeda (1989) and Ariza et al. (2015) avoids  
303 these issues by measuring the capacity of the respiratory ETS on frozen specimens, there are  
304 uncertainties in the choice of ratio to convert from potential respiration to actual  
305 respiration. The inclusion of data collected via both of these methods reduces the influence  
306 of any methodological bias on our results. Additionally, we conduct a bootstrap analysis to  
307 assess uncertainties in our regression model. Although the standard errors calculated for  
308 each coefficient (used to define error bars in Figure 2) were relatively small, they do not  
309 take into account uncertainties in biomass. It is a major challenge, to sample these  
310 mesopelagic fish repeatedly at such a spatial scale, and thus although we are unable to  
311 quantify uncertainties surrounding total biomass estimates at each station, we believe our  
312 analysis is a useful step forward in a complex and poorly-studied area.

313 To allow comparison to studies compiling larger data sets of fish metabolism, we reran our  
314 regression model using the same data but with respiration rates in units of  $\mu\text{L O}_2 \text{ Ind.}^{-1} \text{ h}^{-1}$

315 ( $R_{IND}$ ), rather than mass specific respiration. This allows us to calculate the mass scaling  
316 coefficient ( $a_1$ ) to compare with other studies.

317

$$318 \quad \ln(R_{IND}) = a_0 + a_1 \times \ln(WM) + a_2 \times T \quad (5)$$

319

320 This reveals a mass scaling coefficient of 0.734 (0.682-0.785), comparing well to the  
321 coefficients found by Ikeda (2016) (0.843-0.925), Clarke and Johnston (1999) (0.79-0.83) and  
322 Winberg (1956) (0.687-0.930). This gives confidence that the myctophid respiration dataset  
323 is sufficient to capture relationships between respiration, mass and temperature. Our  
324 compiled data set covers several orders of magnitude in WM (0.026 – 19.2 g), and a wide  
325 temperature range from 0.5 to 27 °C. However, these studies in themselves are subject to  
326 limitations as discussed above.

327 Although relations between mass and metabolic rate have been found when examining  
328 organisms over many orders of magnitude in size (Brown et al. 2004), there is much scatter  
329 around this relationship. A review by Seibel and Drazen (2007) highlighted a 300-fold  
330 variation in metabolic rates between the fastest and slowest marine animals that was  
331 independent of body mass and temperature. Potential differences in locomotory capacity  
332 between the myctophid species used to develop our regression model and those sampled in  
333 our study region therefore adds to the uncertainty in our calculated respiration rates.

334 *Species contribution to total respiration*

335 We find that, for the three sites analysed here, NSS, MSS, and SSS, *Electrona carlsbergi*,  
336 *Electrona antarctica* and *Gymnoscopelus braueri* were the dominant contributors to  
337 respiration. These species were also dominant in terms of total biomass (Collins et al. 2012,  
338 Saunders et al. 2014, 2015a) highlighting that, of the terms in our regression model, total  
339 biomass is a more important determinant of community respiration than individual fish  
340 mass or temperature when considering the Scotia Sea as a whole. This is likely because the  
341 range in temperatures across our study site is small (-1.46 – 3.31 °C). However, the  
342 differences in species composition regionally within the Scotia Sea likely contributes to the  
343 regional differences we see in total respiration (Figure 2).

344 Collins et al. (2012) noted a higher species diversity in the northern Scotia Sea where  
345 temperatures are warmer. This is likely related to the need to attain a greater body size at  
346 the colder temperatures of the southern Scotia Sea, hence preventing smaller species and  
347 intra-specific life stages from penetrating the southernmost regions (Saunders & Tarling  
348 2018). Similar macroecological trends in diversity and body size have also been reported for  
349 fish communities globally (Fisher et al. 2010a, b). *G. braueri* and *E. antarctica* were the  
350 dominant species in terms of abundance in the southern Scotia Sea, whereas *E. carlsbergi*,  
351 *Krefftichthys anderssoni* and *Protomyctophum bolini* were dominant in the northern stations  
352 (Saunders et al. 2014, 2015a, b). The size of *G. braueri* and *E. antarctica* (34-162 mm and 24-  
353 115 mm SL respectively) is larger than that of *E. carlsbergi*, *K. anderssoni* and *P. bolini* (68-90  
354 mm, 15-74 mm and 23-66 mm SL respectively), which may in part explain why total  
355 respiration rates in the SSS were typically higher than those in the NSS.

356 Additionally, as species-specific respiration rates are calculated from a general regression  
357 for myctophids, if there are large inter-species variations in respiration (e.g. due to

358 differences in locomotory capacity, diet and behaviour etc), it is possible that less abundant  
359 species could make greater contributions to total respiration than we have estimated here.  
360 However, there are currently insufficient data to develop species-specific mass-respiration  
361 relationships. It is difficult to collect healthy, live fish from mesopelagic depths for use in  
362 incubation experiments, and *in situ* incubations at depth are not yet feasible for the majority  
363 of scientific research cruises. We suggest that estimating respiration through the  
364 measurement of ETS activity (Packard & Christensen 2004, Ariza et al. 2015) provides a good  
365 alternative, particularly in revealing interspecific differences.

#### 366 *Seasonal patterns in total respiration*

367 Comparison of integrated respiration at NSS, MSS, and SSS (Figure 2) highlights strong  
368 seasonality in the NSS compared to MSS and SSS. As *E. carlsbergi*, a predominantly copepod  
369 feeding species (Saunders et al. 2015a), accounted for most of the biomass and myctophid  
370 respiration at the NSS site in spring, it is possible that high respiration here was driven by  
371 the large phytoplankton blooms (Korb et al. 2008, 2012) and high mesozooplankton  
372 abundances (Ward et al. 2012) that occur in the region. It has been suggested that *E.*  
373 *carlsbergi* may be associated with warm water eddies from the Polar Front (Collins et al.  
374 2012) which, if more prevalent in spring, could explain the seasonal decline in the  
375 contribution of *E. carlsbergi* to myctophid respiration at NSS. The dominance of *E. carlsbergi*  
376 to total respiration at NSS in spring highlights that migration behaviour and oceanic  
377 transport mechanism from more remote regions can be an important factor in community  
378 respiration in the Southern Ocean.

379 Whereas total respiration was greatest in spring at NSS, the maximum respiration occurred  
380 in summer and autumn at MSS and SSS respectively. The spring peak at NSS may be related

381 to the aforementioned migration patterns of *E. carlsbergi*. The later peak in myctophid  
382 respiration in the southern Scotia Sea may be linked to ice cover, with the timing of ice  
383 retreat influencing the development of zooplankton (Korb et al. 2012), which are the prey  
384 for the myctophid species at our study site (Saunders et al. 2014, 2015a). During the same  
385 Discovery 2010 cruises, Ward et al. (2012) observed highest zooplankton abundances in the  
386 autumn in the southern Scotia Sea.

387 It is very interesting that, despite the low temperatures of the SSS station (-1.46 to 0.95 °C,  
388 based on mean net temperatures), total respiration rates are still high and comparable to  
389 both MSS and NSS where temperatures are higher (Figure 2). Thus, despite much higher  
390 zooplankton abundances in the NSS, in terms of myctophid respiration, total respiration is  
391 actually higher in the SSS. The higher abundance of myctophids in the SSS likely explains  
392 these regional patterns in total respiration, with higher abundances perhaps relating to food  
393 availability, or to the refuge from predation that the sea ice zone provides. Krill abundances  
394 are high across the Scotia Sea (Atkinson et al. 2008), but more krill are found in the southern  
395 Scotia Sea (Fielding et al. 2012) where most spawning occurs (Murphy et al. 2007).

396 Therefore, higher abundances of krill in the sea ice zone, particularly of smaller life stages  
397 that fall more within the prey size spectra for myctophids may explain, at least in part, the  
398 higher abundances of some myctophid species in the southern Scotia Sea.

399 Since there are regional differences in prey availability, and myctophids can select larger,  
400 more energy rich copepodite stages when feeding (Shreeve et al. 2009), prey quality may  
401 also play a role in the regional patterns in total respiration. Additionally, as krill typically  
402 have a higher energetic density than copepods (Schaafsma et al. 2018), the increase in krill  
403 predation by *E. antarctica* with increasing latitude southwards (Saunders et al. 2014) could

404 support higher metabolic activities and contribute to higher total respiration at SSS.  
405 However, as our respiration estimates are based primarily on patterns of myctophid  
406 abundance, it would be useful to validate our finding of higher respiration rates in the SSS  
407 by direct measurements of respiration at these sites. If abundance is indeed the primary  
408 driver, then the high spatio-temporal variability in myctophid distribution and abundance  
409 (Collins et al. 2012) has important consequences for active carbon fluxes in the Scotia Sea.

#### 410 *Respiratory carbon flux*

411 We calculate a maximum respiratory carbon flux of 0.05-0.28 mg C m<sup>-2</sup> d<sup>-1</sup> based on net  
412 catch data that has not been corrected for catch efficiency. This is at the low end of previous  
413 estimates of myctophid/micronekton respiration (Table 2) even when rates are adjusted for  
414 differences in *in situ* temperatures. Individual fish WM ranged from 0.03 to 78.34 g (mean  
415 4.38 g) compared to 0.085 to 0.225 g (mean 0.163 g) in the study of Ariza et al. (2015). As  
416 respiration rates are higher for larger individuals, it is surprising that respiratory carbon  
417 fluxes calculated by Ariza et al. (2015) are so high, considering the community of small sized  
418 fish in their study. Size is therefore not the only important factor to consider, and  
419 differences in the locomotory capacity and behaviour of the fish species in the various  
420 studies could also contribute to differences in respiratory carbon fluxes. Hidaka et al. (2001)  
421 and Hudson et al. (2014) do not give individual fish weights to allow size based comparisons.  
422 The different methods of sampling and calculation of respiratory flux in the aforementioned  
423 studies make direct comparisons difficult, but it is clear that our estimates sit in the range of  
424 previous estimates.

425 To assess the potential importance of the respiratory carbon flux of myctophid fishes in the  
426 Scotia Sea we compare our data to the gravitational flux of POC at two sediment traps, P2 at

427 1500 m (at NSS site) and P3 at 2000 m (at GB site) (Manno et al. 2015). Between 2008 and  
428 2010, POC fluxes in November at P2 ranged from 0.6 to 3.2 mg C m<sup>-2</sup> d<sup>-1</sup>, and from 7.1 to  
429 13.1 mg C m<sup>-2</sup> d<sup>-1</sup> at P3 in January (Manno et al. 2015). These compare to a maximum  
430 respiratory carbon flux of 0.28 mg C m<sup>-2</sup> d<sup>-1</sup> at NSS and 0.13 mg C m<sup>-2</sup> d<sup>-1</sup> at GB respectively.  
431 The myctophid respiratory carbon flux alone (i.e. excluding other myctophid driven carbon  
432 fluxes via excretion, mortality and defaecation) is equivalent to 9-47% and 1-2% of the  
433 gravitational POC flux at NSS and GB respectively. These are higher than Hidaka et al. (2001)  
434 and Ariza et al. (2015) measured for euphausiids and decapods in the Canary Islands and  
435 western Equatorial Pacific (euphausiid and decapod respiration were equivalent to up to  
436 1.6% and 1.4% of total POC flux respectively). For comparison, data compiled by Steinberg &  
437 Landry (2017) shows that the respiratory fluxes of zooplankton are typically higher (up to  
438 ~30 mg C m<sup>-2</sup> d<sup>-1</sup>) than our estimates for myctophid fish. However, differences in biomass,  
439 temperature and depth, for example, make it hard to compare values directly. Their study  
440 further revealed a positive trend between percent contribution to POC and respiratory flux,  
441 with zooplankton respiratory fluxes < 2 mg C m<sup>-2</sup> d<sup>-1</sup> corresponding to a contribution to POC  
442 flux of <15%. Despite relatively low total respiratory fluxes in comparison to zooplankton,  
443 our data suggest that the percent contribution can still be high for myctophids.

444 Although our estimate of respiratory carbon flux is a maximum, due to possible day-time net  
445 avoidance, actual active rates of respiration will be higher than the routine respiration rates  
446 calculated here, once physiological processes, such as feeding, swimming activity and  
447 reproductive development have been accounted for. The relationship between the active  
448 metabolic rate (the highest rate of energy expenditure) and the basal or standard metabolic  
449 rate (the minimum energy expenditure required to keep the fish alive) can be as high as 14

450 (Steffensen, John 2005). Johnston et al. (1991) measured the oxygen consumption of the  
451 Antarctic teleost fish, *Notothenia neglecta*, finding that active consumption rates were 4-7  
452 times higher than resting rates. The prior feeding conditions, diet and activity level all affect  
453 respiration, and organisms can adjust their rates of respiration in response to variations in  
454 food supply (Brown et al. 2004). It is therefore not possible to explain all the variation in  
455 respiration rates with T and WM alone, and *in situ* rates of active respiration will be higher  
456 than the routine respiration rates estimated here.

457 Fish also contribute to carbon export via mortality, excretion (dissolved organic carbon) and  
458 the production of faecal pellets, such that the total contribution of myctophids to the  
459 transfer of carbon to depth will be greater than we have estimated here. We also estimate  
460 the gut flux, i.e. the flux of POC in faecal pellets containing non-assimilated food. The energy  
461 budgets of Brett and Groves (1979) give a value of 40% for the percentage of respired  
462 carbon that is defecated. The proportion of defecated carbon that is produced in the deep  
463 ocean will depend on the gut clearance time and duration spent at depth. We  
464 conservatively assume that half of the defecation (i.e. 20% of the respiratory flux based on  
465 Brett and Groves (1979)) occurs at depth, and calculate gut fluxes of  $0.01-0.06 \text{ mg C m}^{-2} \text{ d}^{-1}$   
466 for the migrating myctophids at our case study sites. This increases the active flux to  $0.06-$   
467  $0.34 \text{ mg C m}^{-2} \text{ d}^{-1}$  (total respiratory and gut flux). At NSS and GB, this equates to 10.5-56.0%  
468 and 1.2-2.1% of the gravitational POC flux at NSS and GB respectively. Myctophid fishes can  
469 therefore be an important component of the mesopelagic carbon budget, particularly  
470 considering the vertical migrations they undertake (Pakhomov et al. 1996).

471

472 **Concluding remarks**

473 Our analysis of the literature on myctophid respiration rates, and its application to the  
474 Discovery 2010 survey data, reveals that myctophid respiration could indeed make a  
475 significant contribution to fluxes of carbon to the deep ocean in the Scotia Sea. Our  
476 estimates are based on allometric equations and could be improved through the further  
477 integration of direct, species-specific measurements of myctophid respiration. There is also  
478 a need to assess daytime avoidance, for instance, through comparison with acoustic  
479 observations. Given the extent of their potential contribution, it is now key that future work  
480 further constrains the levels of carbon flux generated by myctophid fish so that they may be  
481 appropriately included in global biogeochemical models.

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#### 488 Data Availability

489 The fish length and weight data utilised in this study can be accessed at the following DOI: XXXXXXXX

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- 636

637 **Tables**638 **Table 1: Data sources for respiration rates of myctophid species**

Reference	Location	Myctophid species	Experimental Temperature (°C)	Wet mass range (g)	Range in species maximum length (cm) ^
Donnelly & Torres (1988)	Eastern Gulf of Mexico	<i>Diaphus mollis</i> , <i>Lampanyctus nobilis</i> , <i>Lepidophanes guentheri</i> , <i>Myctophum affine</i>	7-20	0.112-6.155	6.6-12.4
Torres et al. (1979)	Southern California	<i>Diaphus theta</i> , <i>Lampanyctus regalis</i> , <i>Lampanyctus ritteri</i> , <i>Parvilux ingens</i> , <i>Stenobranchius leucopsaurus</i> , <i>Symbolophorus californiensis</i> , <i>Tarletonbeania crenularis</i> , <i>Triphoturus mexicanus</i>	5-13	0.9-13.7	7.0-21.0
Torres & Somero (1988)	Antarctica	<i>Electrona antarctica</i> , <i>Gymnoscopelus braueri</i> , <i>Gymnoscopelus opisthopterus</i>	0.5	1.0-40.0	11.5-16.2
Ariza et al. (2015)	Canary Islands	<i>Lobianchia dofleini</i>	17.5-19	0.085-0.225	5.0
Ikeda (1989)	Coral Sea, South Pacific	<i>Symbolophorus evermanni</i> , <i>Centrobranchus nigroocellatus</i> , <i>Myctophum spp.</i>	24-27	0.026-1.101	5.0-8.0

639 ^ Maximum lengths (SL, with the exception of species in the study of Torres et al. 1979 which are total length)  
640 of each species have been obtained from Fish Base (Froese & Pauly 2018), and we present here the range in  
641 these lengths for the species within each study.

**Table 2: Comparison of respiratory carbon fluxes ( $\text{mg C m}^{-2} \text{d}^{-1}$ ) calculated in this study, and in the literature.**

Reference	Location	Site	Taxa	Migrant biomass ( $\text{mg C m}^{-2}$ )	Temperature at depth ( $^{\circ}\text{C}$ ) <sup>^</sup>	Respiratory Flux * ( $\text{mg C m}^{-2} \text{d}^{-1}$ )	Respiratory Flux at 2 $^{\circ}\text{C}$ ** ( $\text{mg C m}^{-2} \text{d}^{-1}$ )
This study <sup>a</sup>	Southern Ocean	JR161 WSS	Myctophidae	49.8	2.0	0.05 <sup>+</sup>	0.05 <sup>+</sup>
		JR161 NSS		520.6	2.1	0.28 <sup>+</sup>	0.28 <sup>+</sup>
		JR177 GB		238.5	1.7	0.13 <sup>+</sup>	0.13 <sup>+</sup>
		JR177 MSS		407.1	0.7	0.27 <sup>+</sup>	0.33 <sup>+</sup>
Ariza et al. (2015) <sup>b</sup>	Canary Islands	Time-series station (north of Gran Canaria)	Migratory fish	168	12	2.68	0.69
			Migratory nekton <sup>c</sup>	201	12	2.92	0.7
Hudson et al. (2014) <sup>a</sup>	North Azores	Reykjanes Ridge	Migratory	5.2	6.6	0.005-0.027	0.003-0.014
		Azorean Zone	Myctophidae	40	11.8	0.046-0.271	0.012-0.071
Hidaka et al. (2001) <sup>a</sup>	Western equatorial pacific	Station 15	Migratory Myctophidae	462.5	9.3	1.98	0.73
		Station 16		248.9	9.3	1.06	0.39
		Station 8	Night-time Myctophidae	539.5	9.3	2.31	0.86
		Station 10		406.5	9.3	1.74	0.64
		Station 13		716.92	9.3	3.07	1.1

<sup>^</sup> Temperature depth ranges as follows: This study: mean 400-1000 m, Ariza et al. (2015): approximate temperature 400-500m, Hudson et al. (2014): mean 200-750 m, Hidaka et al. (2001): 400 m

\*Flux below following depths: This study: 200 m, Ariza et al. (2015): 150 m, Hudson et al. (2014): 200 m, Hidaka et al. (2001): 160 m.

\*\* Adjusted to 2 $^{\circ}\text{C}$  based on a Q10 of 3.9 for myctophids (Donnelly & Torres 1988)

<sup>+</sup> Maximum respiratory carbon flux as day-time net catches have not been corrected for capture efficiency

<sup>a</sup> uncorrected for capture efficiency

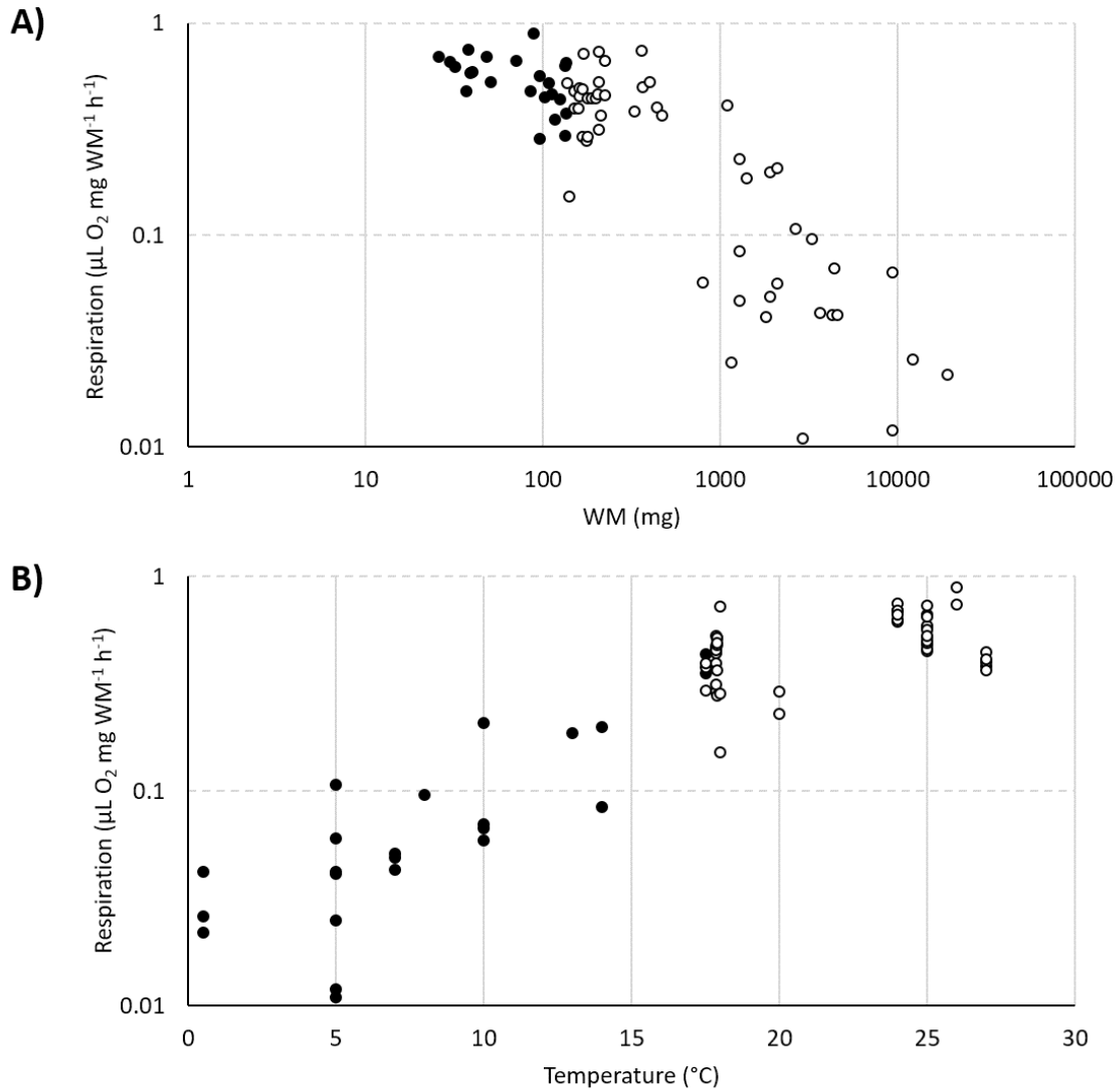
<sup>b</sup> assumes 14% capture efficiency

<sup>c</sup> fish, euphausiids and decapods

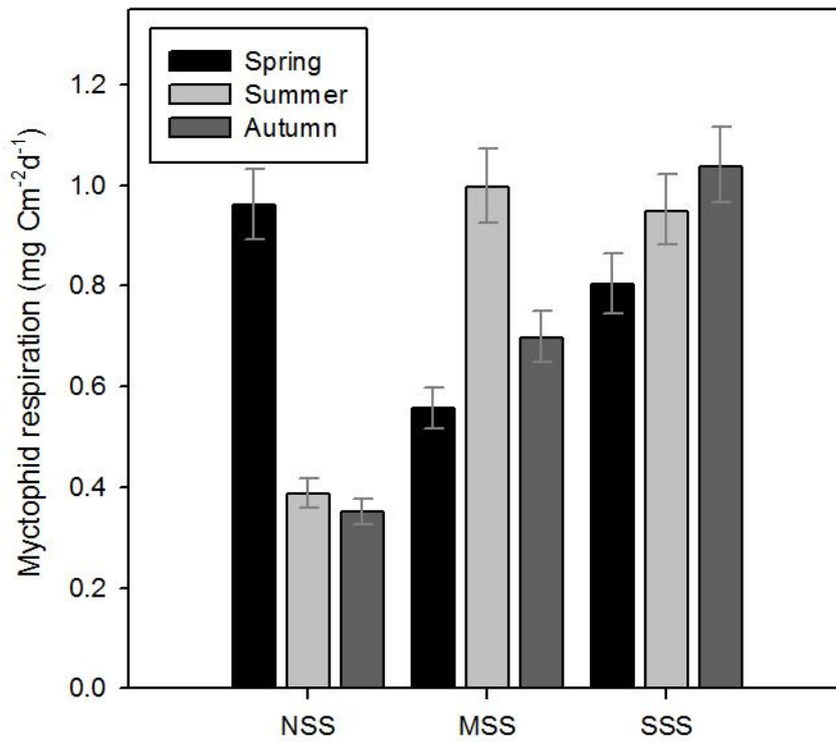
**Table 3: Sensitivity analysis of respiratory carbon flux estimates. Flux estimates have been recalculated based on day-time net capture efficiencies of 14%, 25% and 50%.**

Site	Respiratory flux (mg C m <sup>-2</sup> d <sup>-1</sup> )			
	100%	14%	25%	50%
JR161 WSS	0.05	-0.00	0.02	0.04
JR161 NSS	0.28	0.24	0.26	0.27
JR177 GB	0.13	-0.00	0.06	0.11
JR177 MSS	0.27	0.25	0.26	0.27

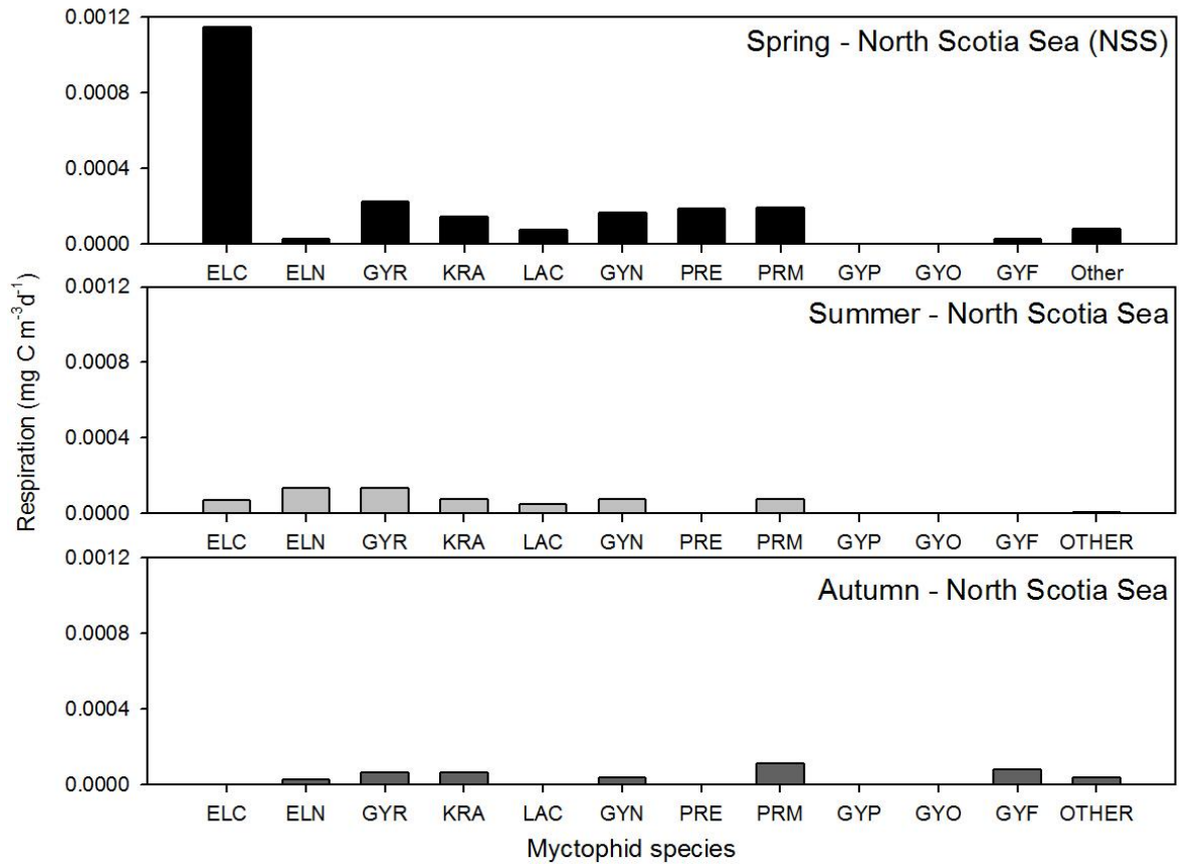
## Figures



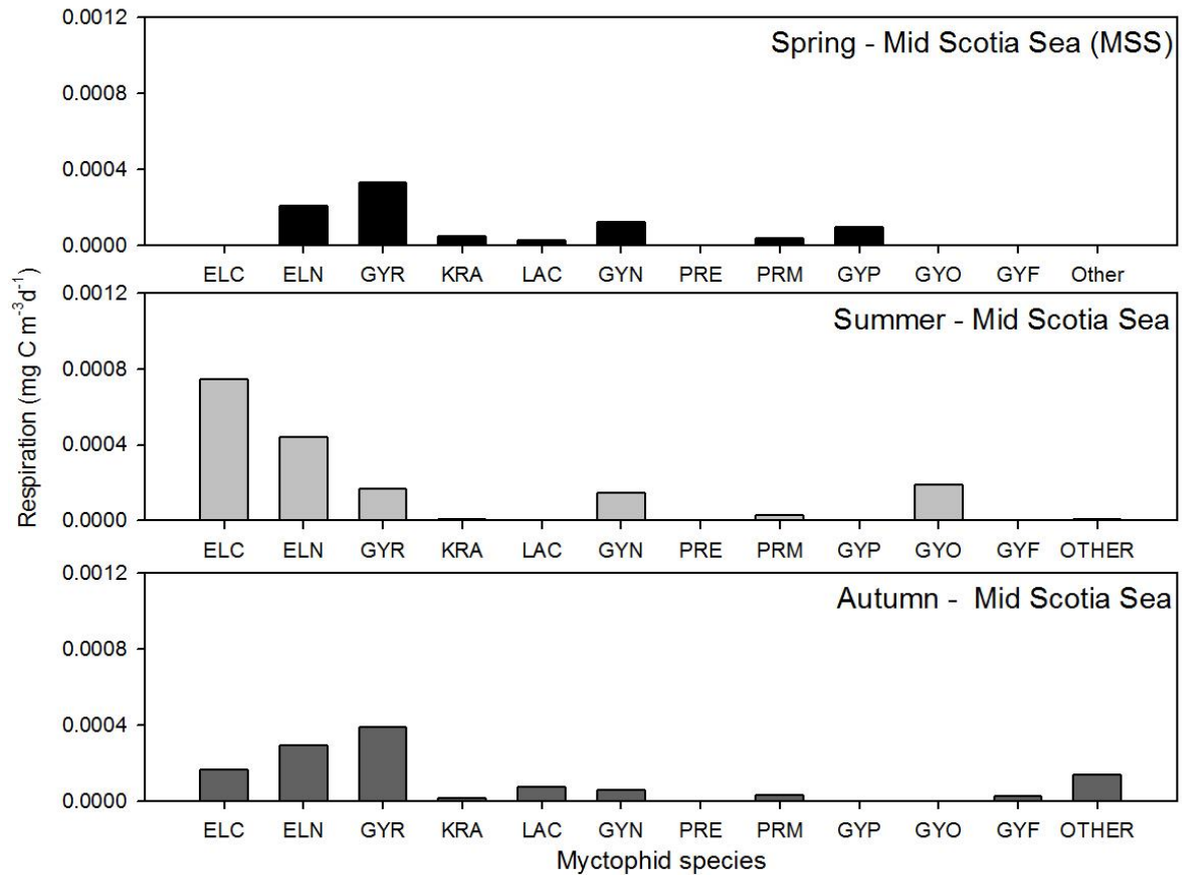
**Figure 1: Literature compilation of respiration rates (mass specific) of myctophid fishes versus A) wet mass (WM), and B) temperature. Note the logarithmic scales. Filled black circles show data from direct oxygen consumption experiments, and open circles show respiration estimated from ETS measurements.**



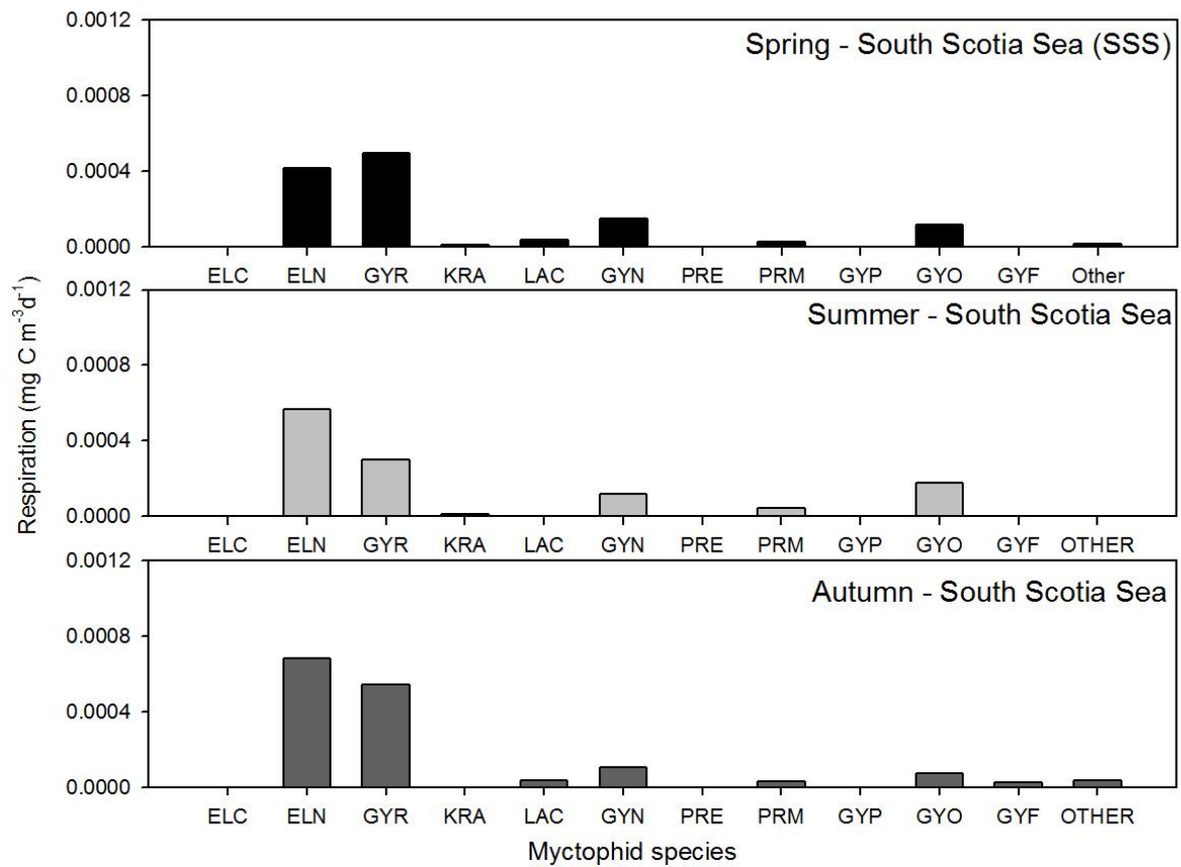
**Figure 2: Seasonal changes in total myctophid respiration ( $\text{mg C m}^{-2} \text{d}^{-1}$ , depth integrated 0-1000 m) in the North Scotia Sea (NSS), Mid Scotia Sea (MSS) and South Scotia Sea (SSS). Data are from night-time hauls only. Error bars display the standard error of bootstrapping analysis (100 runs) of our length-mass regression only (see Methods for full details).**



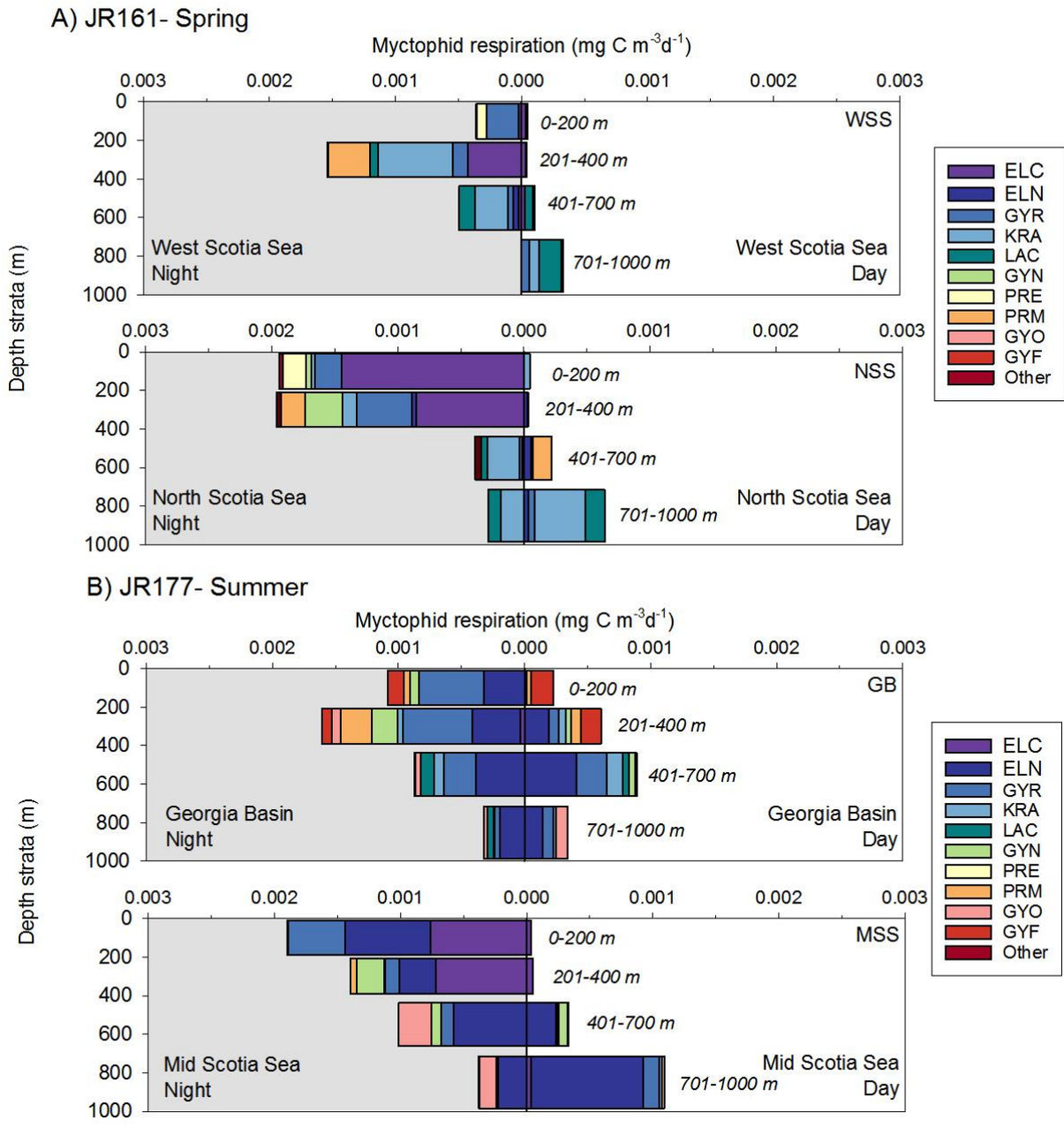
**Figure 3: Seasonal change in total respiration (mg C m<sup>-3</sup> d<sup>-1</sup>) of myctophid fishes caught in the upper 1000 m at the North Scotia Sea (NSS) station. Species code names are as follows: ELC= *Electrona carlsbergi*, ELN = *Electrona antarctica*, GYR= *Gymnoscopelus braueri*, KRA= *Krefftichthys anderssoni*, LAC= *Nannobranchium achirus*, GYN= *Gymnoscopelus nicholsi*, PRE= *Protomyctophum tenisoni*, PRM= *Protomyctophum bolini*, GYP= *Gymnoscopelus piabilis*, GYO= *Gymnoscopelus opisthopterus*, GYF= *Gymnoscopelus fraseri*, OTHER= Other myctophid species. Data from night-time hauls only. Zero values represent species absence.**



**Figure 4: Seasonal change in total respiration (mg C m<sup>-3</sup> d<sup>-1</sup>) of myctophid fishes caught in the upper 1000 m at the Mid Scotia Sea (MSS) station. Species code names are as of Figure 3. Data from night-time hauls only. Zero values represent species absence.**



**Figure 5: Seasonal change in total respiration (mg C m<sup>-3</sup> d<sup>-1</sup>) of myctophid fishes caught in the upper 1000 m at the South Scotia Sea (SSS) station. Species code names are as of Figure 3. Data from night-time hauls only. Zero values represent species absence.**



**Figure 6: Contribution of the dominant myctophid species to the depth stratified respiration in A) spring – cruise JR161 and B) summer – cruise JR177. Total respiration ( $\text{mg C m}^{-3}\text{d}^{-1}$ ) for each species has been calculated for both the day and night (grey shaded graph) net hauls. Species code names are as of Figure 3.**

Supplementary Material

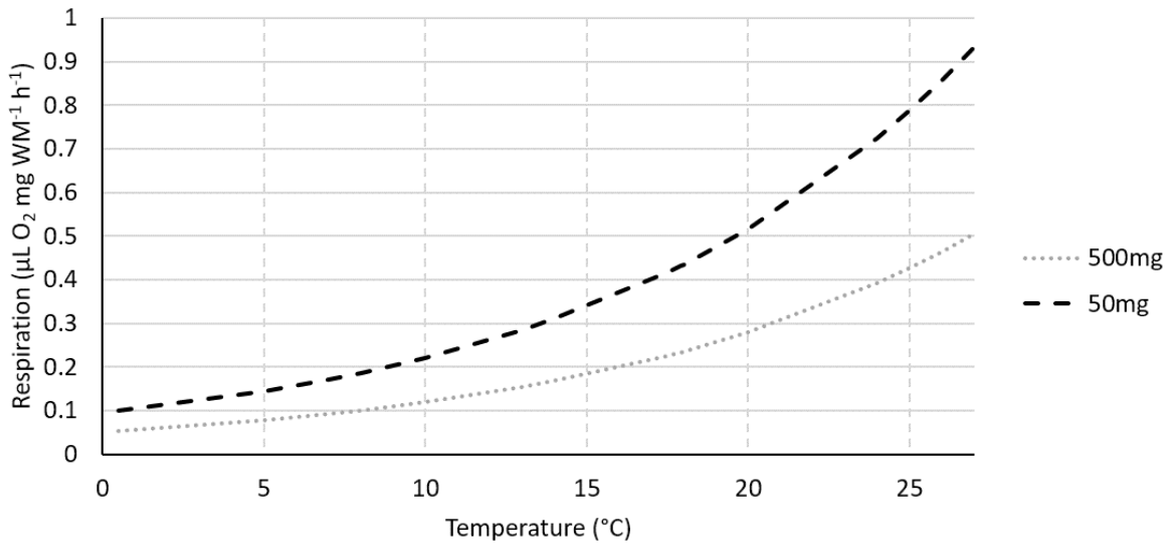


Figure S1: Calculated relationship between temperature (°C) and mass specific respiration rate (µL O<sub>2</sub> mg WM<sup>-1</sup> h<sup>-1</sup>) for myctophid fishes of wet mass 50 mg (black dashed line), and 500 mg (grey dotted line).

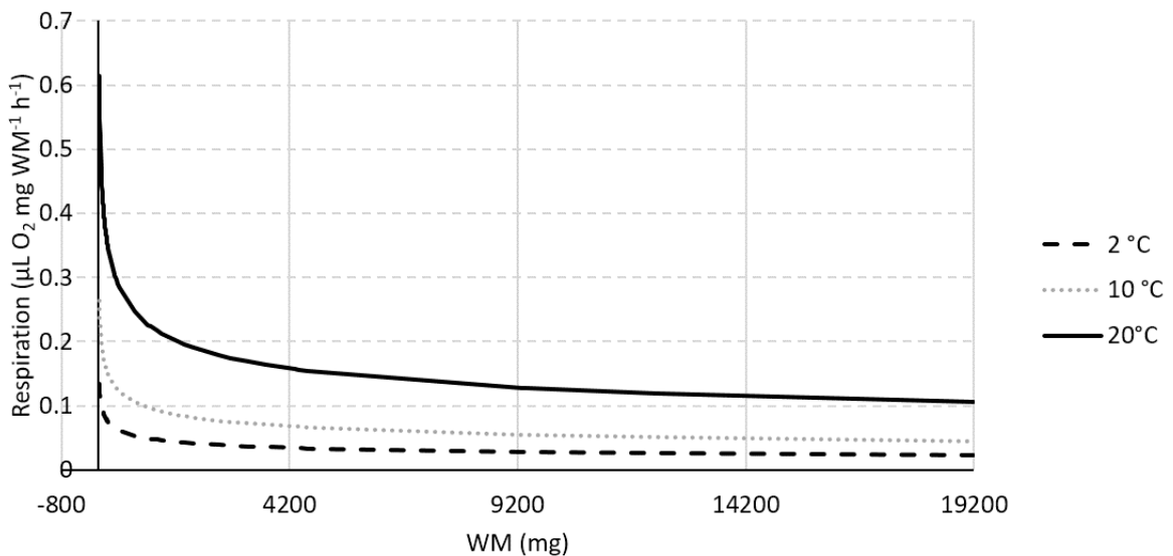


Figure S2: Calculated relationship between temperature (°C) and mass specific respiration rate (µL O<sub>2</sub> mg WM<sup>-1</sup> h<sup>-1</sup>) for myctophid fishes at temperatures of 2 °C (black dashed line), 10 °C (grey dotted line), and 20 °C, solid black line.

**Table S1: Length- Mass regressions used for conversion from standard length (SL, in mm) to wet mass (WM, in g). Lower and upper 95% confidence intervals are also given for each coefficient. Regression:  $WM = a SL^b$**

Species/Genera Name	$a$	Lower	Upper	$b$	Lower	Upper	R <sup>2</sup>
<i>Electrona carlsbergi</i>	$2.09 \times 10^{-05}$	$9.51 \times 10^{-06}$	$4.59 \times 10^{-05}$	2.90	2.72	3.08	0.7214
<i>Electrona antarctica</i>	$3.72 \times 10^{-06}$	$3.22 \times 10^{-06}$	$4.30 \times 10^{-06}$	3.27	3.24	3.31	0.9599
<i>Gymnoscopelus fraseri</i>	$3.53 \times 10^{-06}$	$1.31 \times 10^{-06}$	$9.51 \times 10^{-06}$	3.24	3.00	3.47	0.8811
<i>Gymnoscopelus nicholsi</i>	$2.87 \times 10^{-06}$	$2.02 \times 10^{-06}$	$4.08 \times 10^{-06}$	3.25	3.18	3.33	0.9936
<i>Gymnoscopelus braueri</i>	$4.58 \times 10^{-06}$	$3.60 \times 10^{-06}$	$5.82 \times 10^{-06}$	3.11	3.06	3.17	0.9326
<i>Krefflichthys anderssoni</i>	$9.05 \times 10^{-06}$	$7.49 \times 10^{-06}$	$1.09 \times 10^{-05}$	3.02	2.97	3.07	0.9599
<i>Nannobrachium achirus</i>	$8.14 \times 10^{-06}$	$5.17 \times 10^{-07}$	$1.28 \times 10^{-02}$	2.49	1.45	3.54	0.4259
<i>Protomyctophum tenisoni</i>	$1.39 \times 10^{-05}$	$9.74 \times 10^{-06}$	$1.97 \times 10^{-05}$	2.94	2.84	3.03	0.9589
<i>Protomyctophum bolini</i>	$1.98 \times 10^{-05}$	$1.34 \times 10^{-05}$	$2.92 \times 10^{-05}$	2.88	2.77	2.98	0.8926
<i>Protomyctophum choriodon</i>	$1.27 \times 10^{-05}$	$3.24 \times 10^{-06}$	$4.94 \times 10^{-05}$	2.98	2.66	3.30	0.8779
<i>Gymnoscopelus opisthopterus</i>	$1.20 \times 10^{-06}$	$3.36 \times 10^{-07}$	$4.25 \times 10^{-06}$	3.43	3.16	3.71	0.9874
<i>Electrona</i>	$3.26 \times 10^{-06}$	$2.84 \times 10^{-06}$	$3.74 \times 10^{-06}$	3.31	3.28	3.34	0.9563
<i>Gymnoscopelus</i>	$4.49 \times 10^{-06}$	$3.61 \times 10^{-06}$	$5.59 \times 10^{-06}$	3.12	3.07	3.17	0.9351
<i>Protomyctophum</i>	$1.24 \times 10^{-05}$	$9.89 \times 10^{-06}$	$1.55 \times 10^{-05}$	2.99	2.93	3.05	0.9453